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APPLICATION NUMBER: 60/590,987

FILING DATE: *July 26, 2004*

RELATED PCT APPLICATION NUMBER: PCT/US04/43969



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072604

PROVISIONAL APPLICATION COVER SHEET

This is a request for filing a PROVISIONAL APPLICATION Under 37 CFR 1.53 (b)(2).

Attorney Docket No.

624.P

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INVENTOR(s)/APPLICANT(s)

LAST NAME	FIRST NAME	MIDDLE INITIAL	RESIDENCE (CITY AND EITHER STATE OR FOREIGN COUNTRY)
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TITLE OF THE INVENTION (280 characters max)

HPV INHIBITORS

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ENCLOSED APPLICATION PARTS (check all that apply)

<input checked="" type="checkbox"/> Specification	Number of pages <u>19</u>	<input type="checkbox"/> Small Entity Statement
<input type="checkbox"/> Drawing(s)	Number of sheets	<input type="checkbox"/> Other (specify) _____

METHOD OF PAYMENT (check one)

<input checked="" type="checkbox"/> The Commissioner is hereby authorized to charge filing fees (as well as any additional fees which may be required by this paper) and credit Deposit Account Number <u>07-1250</u> .	Provisional Filing Fee Amount (\$) \$ 160.00
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The invention was made by an agency of the United States Government of under a contract with an agency of the United States Government.

☒ No.

☐ Yes, the name of the U.S. Government Agency and the Government contract number are:

Respectfully submitted,

SIGNATURE

James J. WongDATE July 26, 2004

TYPED or PRINTED NAME

James J. WongREGISTRATION NO. 34,949
(if appropriate)

☐ Additional inventors are being named on separately numbered sheets attached hereto

2386 U.S. PTO
60/590987

072604

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of: Jianying Wang

For: HPV Inhibitors

Commissioner for Patents

P.O. Box 1450

Alexandria, VA 22313-1450

PROVISIONAL APPLICATION COVER SHEET
(37 C.F.R. § 1.51 (2) (i))

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HPV INHIBITORS

20

Most Prevalent Serious HPV-Mediated Diseases (US and EU)

- **Anal dysplasia in HIV patients**
 - low & high-grade dysplasias ~ 575,000
- **Cervical dysplasia**
 - high-grade lesions ~ 470,000
 - low-grade ~ 12MM
- **Genital warts**
 - ~2.3MM
 - 280,000 patients/yr receive topical pharmaceutical therapies
- **Anal dysplasia in homosexual males**
 - low & high-grade dysplasias ~ 700,000

Why is Gilead Interested in HPV

- Series of nucleotides have in vitro activity in HPV+ cell lines
- Cidofovir and analogs effective in animal models
 - SiHa xenograft and CRPV models
- Cidofovir has shown some efficacy in HPV associated human diseases
 - anogenital warts, cervical intraepithelial neoplasia, respiratory papillomatosis
- HPV-associated diseases are an unmet medical need
- Some indications, particularly anal dysplasia in HIV patients, are a good corporate fit

Human Papillomaviruses

- Small, non-enveloped, DNA virus
- Dependent on host cell for replication
- Epithelial tropism
 - site of occurrence (cutaneous, mucosal)
- Species specific with > 100 human subtypes
 - low and high risk subtypes

HPV Genotypes

- High risk:
 - HPV-16, 18, 31, 33, 35, 45
 - potential to induce malignant proliferation
 - HPV-16,18 responsible for 50-80% of dysplasias
 - untreated dysplasias may develop into cancer
- Low risk:
 - HPV-6, 11
 - responsible for nearly 90% of genital warts

Target Product Profile Overview

- HPV Activity
 - activity against HPV-16,18; ideally activity against HPV-6,11
- Selectivity
 - good selectivity between HPV-infected and non-infected tissue
- Safety
 - minimally irritating to mucosal tissue
 - non-mutagenic
 - ideally, non-teratogenic
- Dosing
 - once-daily dosing acceptable for anal and cervical dysplasias and for genital warts
- Formulation
 - topical gel/foam/cream

Goals for Program

- Topical prodrugs
 - improve potency
 - allow for skin penetration
 - reduce toxicity
- Selectivity in vitro
 - EC₅₀ “normal” cell line/EC₅₀ HPV+ cell line
- Topical efficacy in representative animal models
- Compound with minimal irritation and minimal/no genotoxicity

Key Challenges

- Selectivity index is low in vitro
- Mechanism of action is not understood
- The parent molecules are potentially toxic
 - renal toxicity, mutagenicity, carcinogenicity, local irritation

Screening cascade I; early stage

All compounds

Anti-proliferation
assays

Mechanism of
action studies

Log D, stability,
Solubility, cleavage

7-day EC₅₀ in
SiHa(+) and HEL(-)



Active compounds
(< 10 nM)

7-day EC₅₀ in
MS751(+) and
PHK(-)



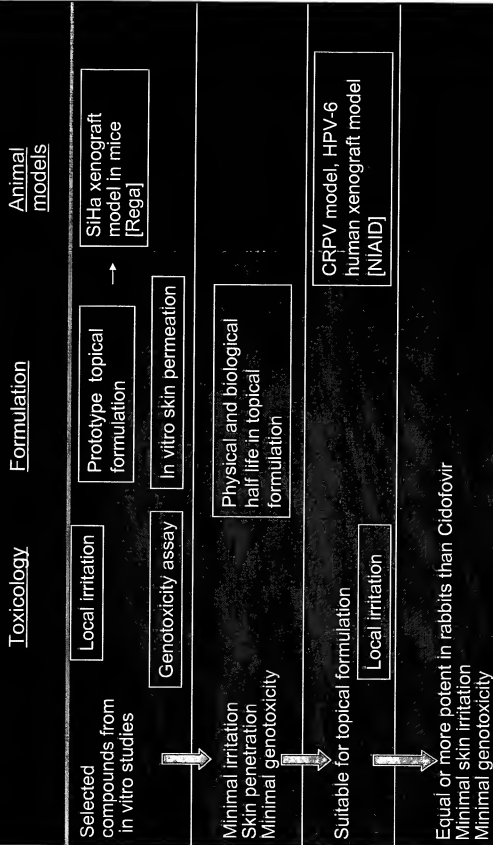
7-day EC₅₀ in
additional cell types

Selected
compounds*

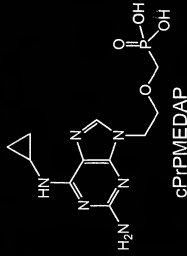
Necrosis (LDH assay)
Apoptosis (Caspase 3)
DNA synthesis (BrdU)
Cell cycle analysis
Metabolic pathway
Mitochondrial toxicity?
Telomerase?
DNA polymerase
assay?

*Selection criteria: EC₅₀, selectivity, stability, etc.

Screening cascade II; late stage



cPrPMEDAP: Scaffold for Prodrug Design



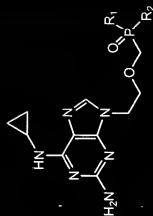
- Anti-proliferative activity against HPV(+) cell lines (EC₅₀ at sub- μ M range)
- Selectivity when compared to HPV(-) cell lines or primary human keratinocytes

Disadvantages of Other Parent Scaffolds

- Cidofovir (HPMPC) has moderate activity but less selective
- PME_A is not active
- PMEDAP and its other analogs are less active
- PMEG has similar activity; synthetically more challenging

Improved Potency of cPrPMEDAP Prodrugs

GS#	R1	R2	EC ₅₀ (nM) in HPV16+ SiHa cell line
8369	OH	OH	284
17429	O- <i>i</i> Pr	O- <i>i</i> Pr	2267
327353	Ala- <i>i</i> Pr	Ala- <i>i</i> Pr	2.5
327238	Ala- <i>i</i> Pr	Ala- <i>i</i> Pr	1.3
327319	Aba-Et	Aba-Et	3.2
327261	Aba-Bu	Aba-Bu	0.20
327352	O ^t Ph	Ala- <i>i</i> Pr	0.50
327383	O ^t Ph	Aba-Bu	0.13
56884	O ^t Ph	Phe-Et	0.60



8369 (cPrPMEDAP)

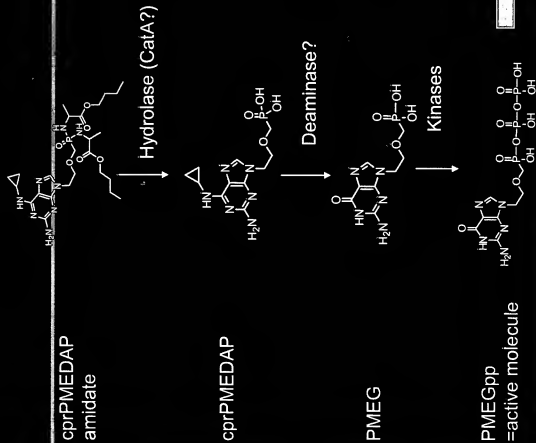
Selection Criteria for Initial *in vivo* Screens

- Key criteria
 - Selectivity HPV+/HPV- cell lines
 - Log D between 1.5 and 2.5
 - Stability in formulation vehicle
- Other
 - Potency in HPV+ cell lines
 - Solubility
 - Cleavage by CAT A

cPrPMEDAP Prodrugs with Good Selectivity

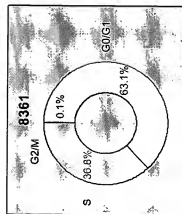
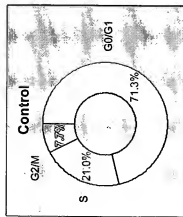
GS#	Structure		Selectivity	
	R1	R2	HEL/SiHa	PHK/SiHa
8369	OH	OH	17	13
327353	Ala-Pr	Ala-Pr	210	31
327238	Ala-IPr	Ala-IPr	559	75
327319	Aba-Et	Aba-Et	135	12
327261	Aba-Bu	Aba-Bu	115	4
327352	OPh	Ala-Pr	164	10
327383	OPh	Aba-Bu	92	22
56864	OPh	Phe-Et	72	58
Podofilox	Active ingredient of condylox		<0.9	0.1
8358	PMEG bis Ala-Bu		11	1.8
AraC	C-analog DNA pol inh		0.11	0.57
Cladribine	A-analog DNA pol inh		nd	1

Mechanism of Selectivity (hypothesis)

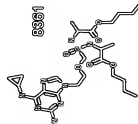
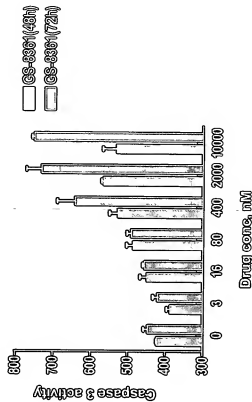


A prototype prodrug arrests SiHa cells at the S-phase of the cell cycle and induces apoptosis

Cell cycle analysis
48 hr



Induction of Apoptosis



Mechanism of Selectivity (experimental plan)

- Compare the rate of metabolic conversion in cells to identify the rate-limiting reaction
 - SiHa (HPV16, sensitive to cprPMEDAP amidates)
 - CaSki (HPV16, resistant)
 - Primary keratinocytes (HPV neg, somewhat resistant)
 - Primary fibroblasts (HPV neg, resistant)

PHK-SiHa Raft Co-cultures (after 10 days of differentiation)



Control



cPTMEDAP 5 µg/mL

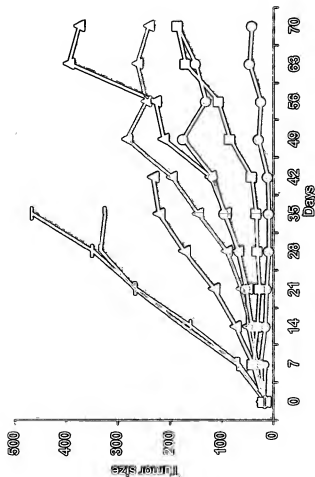


cPTMEDAP 0.5 µg/mL



cPTMEDAP 0.05 µg/mL

SiHa Tumor Model - Athymic Mice Monoamidates (Ala-iPr)



3 weeks treatment, 5 days/week
100 μ l intratumoral

